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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)							
(51) International Patent Classification ⁶ :		(11) International Publication Number: WO 99/35082					
C01B 3/38	A1	(43) International Publication Date: 15 July 1999 (15.07.99)					
(21) International Application Number: PCT/US9	99/006	(81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD,					
(22) International Filing Date: 11 January 1999 (1	1.01.9	KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK,					
(30) Priority Data: 09/005,676 12 January 1998 (12.01.98)	τ	MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM),					

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European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

Published

With international search report.

(54) Title: CONTROL OF HYDROGEN AND CARBON MONOXIDE PRODUCED IN PARTIAL OXIDATION PROCESS

(57) Abstract

A process for enhancing H2 or CO production in a partial oxidation reaction by feeding H2O or CO2 with the feed hydrocarbon and oxygen over a transition metal monolith catalyst such as unsupported Ni monolith or alternatively contacting the hydrocarbon/oxygen first with a noble metal then with a transition metal with the H2O or CO2 being added before or after the noble metal catalyst. The addition of H₂O suppresses CO and enhances H₂ production and the addition of CO₂ suppresses H₂ and enhances CO production. Little steam or CO₂ reforming occurs with the addition of up to 32 % H₂O or CO₂ respectively. Thus, the ratio of H₂:CO which is about 2 in a conventional partial oxidation is manipulated by the addition of either water or CO₂ to the partial oxidation.

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CONTROL OF HYDROGEN AND CARBON MONOXIDE PRODUCED IN PARTIAL OXIDATION PROCESS

BACKGROUND OF THE INVENTION

5 Field of the Invention

The present invention relates to a process and apparatus for production of H₂ or CO by the partial oxidation of hydrocarbons, preferably methane. In particular the partial oxidation is carried out in the presence of water or carbon dioxide under water-gas shift reaction conditions. The invention was made with government support under DOE Grant No. DE-FGO2-88ER13878-AO2. The government has certain rights in the invention.

Catalytic steam reforming of methane is currently the main industrial process to produce synthesis gas (CO and $\rm H_2$). The steam reforming reaction may be represented as:

$$CH_4 + H_2O \longrightarrow CO + 3H_2$$

Reforming is highly endothermic, requiring energy input and also requiring contact times on the order of seconds. The resultant high $\rm H_2/CO$ ratio is also unsuitable for methanol and Fischer-Tropsch synthesis. Therefore additional downstream reactors are usually required to adjust this ratio by water-gas shift reaction at contact time of ≈ 1 second as:

$$CO + H_2O \longleftrightarrow H_2 + CO_2$$

Partial oxidation, on the other hand, is an exothermic reaction which can be represented by the reaction of methane with oxygen as follows:

$$CH_4 + {}^{1}O_2 - \longrightarrow CO + 2H_2$$

To produce synthesis gas by steam reforming, high temperature heat input is primarily required at two process steps. First, sufficient steam at a high temperature and high pressure must be generated for mixing with the hydrocarbon feedstock and, second, the steam reforming of the steam and hydrocarbon mixture must take place at relatively high temperatures and pressures through a bed of solid catalyst. The equipment needed for these two heat transfers at high temperature and high pressure is

necessarily quite expensive. The equipment for the steam reforming step is also costly because it must be adapted to permit the changing of the solid catalyst when that catalyst is spent or poisoned. Heat sources appropriate for the above two process steps are typically provided by fired heaters at high, continuing utility costs, also with high fluegas NOX production consequential to the high temperatures required in the furnace firebox.

The production of synthesis gas by partial oxidation is considered as desirable alternative to steam reforming since it overcomes some of the problems of steam reforming, see for example PCT publication W0 90/06282 90/06297. U.S. Pat. No. 4,844,837 to Heck et al discloses de catalytic partial oxidation method for methane using a monolith catalyst with platinum-palladium, palladiumrhodium, or platinum-rhodium coatings. o U.S. Pat. No. 4,087,259 to Fujitani et al describes a monolith catalyst with a rhodium coating to perform catalytic partial oxidation on gasoline and heavier petroleum fractions. U.S. Pat. No. 5,648,582 to Schmidt et al discloses the partial oxidation of methane at short residence times using metal deposited on a ceramic monolithe espectation

The these conventional partial oxidations, methane daiw reet ' (natural gas) is converted to high purity H2 and CO with a mole ratio of H2:CO 22:0, which is the desired feed ratio for methanol and Fischer-Tropsch plants However, many other applications require different ratios of H2:CO. present synthesis gas shift reactor modifies the prior processes to obtain either high purity H2 or CO.

30 Recent advancements in fuel cell technology have spurred an interest in converting natural gas into hydrogen. Pure hydrogen streams can be produced by steam reforming followed by high temperature shift using an Fe based catalyst, and low temperature shift using a Cu based catalyst. For natural gas to be an effective H2 source for fuel cells, the present natural gas conversion technology must be simplified, preferably to a single, highly selective small catalytic reactor. Other applications

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include acetic acid production, which requires pure CO feeds. We have discovered that the H₂:CO product ratio can be altered by the addition of CO₂ or H₂O through the watergas shift reaction while the conversion of CH₄ remains constant, indicating negligible reforming is occurring.

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partial oxidation of hydrocarbons such as methane (natural gas) by contacting a feed containing the hydrocarbon and oxygen and H2O or CO2 through a catalyst zone containing a catalystically effective amount of at least one transition metal monolith catalyst under partial oxidation conditions. A preferred monolith catalyst is a nickel metal monolith. When H2O is feed, the product shifts toward the H2 and when CO2 is present the product shifts toward CO, thus the presence of either water or CO2 provides the means to adjust the H2:CO ratio, preferably in the range of more than about 2 to about 6 when water is added and in the range of less than about 2 to about 0.5 when CO2 is added.

20 It appears that the presence of water or CO2 in the reaction zone according to the present invention produces a

water-gas shift rather than reforming coupe in a further embodiment the process comprises a first contacting a feed comprising methane and oxygen feed with 25 or without H2O or CO2 with a noble metal coated onto a metal monolith or between the noble metal monolith and the transition metal monolith.

Because partial oxidation reactions are exothermic, it is not necessary to add external heat to the system other operation to obtain ignition of the catalyst.

BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1 shows the conversion of CH_A and selectivities to

35338 Fig. 2 shows the conversion, of CH4 and selectivities to

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Fig. 4 shows the conversion of CH₄ and selectivities to

Fig. 5 shows the conversion of CH_4 and selectivities to co with $H_2\theta$ addition over Rh.

Fig. 6 shows the conversion of CH_4 and selectivities to CO with $H_2^{\circ}O$ addition over Pt.

Fig. 7 compares the selectivity to CO in the product with the addition of H2O over Ni, Rh, and Pt.

Fig. 6 compares the ratio of the $\rm H_2$ in the product to the $\rm CH_4$ in the feed with the addition of $\rm H_2O$ over Ni, Rh, and Pt.

Fig. 9 compares the ratio of CO/CO2 in the product with the addition of H2O over Ni, Rh and Pt.

Fig. 10 compares the conversion of CH_4 with the addition of $H_2\bar{O}$ over Ni, Rh, and Pt.

Fig. 11 discloses a single bed catalyst reaction system according to the present invention.

Fig. 12 discloses a multibed catalyst reaction system according to the present invention.

Fig. 13 discloses an alternative multibed catalyst reaction system according to the present invention.

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The present catalyst is comprised of a monolith structure either composed of the metal of the catalyst or coated with the metal of the catalyst. The metals may be in the oxide form during use in the present process. The thickness of the monolith through which the feed gas mixture must pass is from 1 mm to 2 cm. Catalyst contact time ranges from 0.1 to 20 milliseconds when using a monolith of 50% porosity and 0.2 to 1 cm in depth. Under operating conditions, this corresponds to Gas Hourly Space Velocity (GHSV) of 60,000 to 3,000,000 hr 1

Ceramic foam monoliths have been found in the present invention to create the superior mass transfer characteristics necessary if high space velocities are to be used. The metal monolith may be prepared as metal foam or sintered particles of metal. The solid metal monoliths exhibit superior heat transfer properties but may require

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substantial amounts of very expensive metals in some embodiments. Thus, in some applications the metal coated ceramics will be the catalyst of choice. As used herein the term "metal monolith" shall include both the solid metal monoliths and the metal coated ceramic monoliths. The solid metal monoliths may be produced by any method, for example foaming, sintering and fusing.

The reactor is started from ambient temperature through at some otherwise of a mixture of light hydrocarbons or ammonia and mair preheated to about 200°C and then introduced to the monolith catalyst at an appropriate temperature at which combustion will occur. After combustion has established a monolith catalyst temperature of near 1000°C, preheat and ammonia is stopped. The feed gas mixture of hydrocarbons (methane) and oxygen is then fed to the catalyst zone at a temperature of from 25° to 450°C. Thus, the gas feed mixture of the present invention does not require M. approprietating to, near- its ignition temperature prior to introduction to the catalyst, thereby avoiding the 20 production of 502 and H20 and the concurrent reduction of of the selectivity for H2 and CO. With the introduction of water or carbon dioxide to the reaction in accordance with the present invention, the preheat of the feed is adjusted an inclination maintain the established outlet temperature. sed w.25 wiscarther amounts so of many hydrocarbon, at H2O, CO2, and oxygen and landintroduced into the partial oxidation (catalyst zone) are controlled to provide 02: Caratios of from about, 0.2 to 0.8 doubted or and H2O or CO2; C ratios of about 0.5 to 5... The process is a prime mearried out from about atmospheric pressure to about 2000 1941.130 .dr.psig. The amount of water or CO2 added is preferably up to word > (12:45% more preferably 5 to 35% of the gaseous feed.

Because the hydrocarbon conversion remains constant with or without the H₂O or CO₂ it appears that little or no reforming is occurring. Thus the present partial oxidation in the presence of H₂O or CO₂ is characterized as occurring in the substantial absence of reforming. The predominant reaction in the presence of water or CO₂ is characterized as a water-gas shift reaction.

in a preferred embodiment of the present invention, catalytic partial oxidation, an exothermic reaction, takes place in a catalyst monolith having a specified transition metal alone or preceded by a noble metal dispersed to produce a gas which is rich in carbon monoxide and hydrogen. The reaction in the catalytic partial oxidation zone is exothermic and the zone is therefore also referred to as an exothermic catalyst zone. The exothermic, catalytic partial oxidation zone comprises either solid 10 metal monolith and/or a monolithic catalyst carrier or carriers on which transition or transition and noble metal catalyst is dispersed. Such catalyst can effectively catalyze the partial oxidation of in addition to, gaseous and lighter hydrocarbon liquids such as natural gas or paraffinic naphtha, heavier hydrocarbon liquids such as diesel oil, number 2 fuel oil, and coal derived liquids. As compared to a non-catalytic combustion process such as conventional, non-catalytic partial oxidation, catalytic partial oxidation as described above enables the 20 utilization of lesser amounts of oxygen and lower temperature levels to both oxidize a portion of the feed and crack heavier feedstocks to lighter hydrocarbon fractions while raising the temperature of the reactant mass for subsequent treatments: 2 Generally, at least about 25 25 half the hydrocarbon feed stock is partially oxidized in the catalytic partial oxidation zone to produce primarily carbon monoxide and hydrogen and heat. Substantially all of to ship of the oxygen introduced into the catalytic partial oxidation The oxygen introduced into the catalytic partial exidation zone is Consumed in the catalytic spartial coxidation astep. sols tales oxygen may be provided by any suitable loxygen-containing oxidant gas with ich termois used in the claims to include Tair air enriched with oxygen; oxygen; oxygen mixed with 16.18354 Wother gases. The seffluent gas from the catalytic partial oxidation zone contains primarily CO, H2, H2O, N2, C2 to C4 and other flighter hydrocarbons, including olefins, and, depending upon the sulfur content of the feedstock, H2S and

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Methane is the preferred feed for ${\rm H}_2$ cos. production. Mag to Posts of

The presence of H20 (steam) results in a favorable shift in the product of the H2:COC ratio to H2; whereas the 5 presence of CO2 shifts the ratio in favor of CO. .

The combination of features provided by the present invention provides a highly efficient and flexible method number of converting various typestof hydrocarbonaceous feeds to a hydrogen-rich gas. For example, the combination of 10 1 features provided by the process of the present invention add of provides a highly efficient process of manufacturing a various types of by converting is deport synthesis gas mydrocarbonaceous feeds, including hydrocarbon feeds, to a to be mitrogen and hydrogen-rich gas suitable for use in ammonia synthesis: By utilizing the catalytic partial oxidation process as described, a wide variety of hydrocarbonaceous feeds may be efficiently and economically converted into a Services a hydrogen-rich agas. The late to the late to

The Monolithic Partial Oxidation Catalysts The partial oxidation catalyst either comprises or is

supported on a monolithic carrier, that is, a carrier of Little type comprising one or more monolithic bodies having a Andreser epiurality of ofinely divided gas flow passages extending Jacks therethrough and Suchamonolithic carrier members are soften as basis referred to as "honeycomb" type carriers and are well known villasming and the art. and preferred form of such carrier is made of a to listy direfractory, substantially inert, rigid material which is capable of maintaining its shape and a sufficient degree of medianicalistrength at high temperatures, for example, up ab : 30 Typically, a material is . The teselected of the support which exhibits a low thermal characters of coefficients of expansion, egood, thermal shock resistance obsides and, though note always, whow thermal, conductivity. The part of general atypes of material for construction of 35 carriers are known. One is a ceramic-like porous material comprised of one sor more metals oxides, for example, alumina, alumina-silica, alumina-silica-titania, mullite,

cordierite, zirconia, zirconia-spinal, zirconia-mullite,

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silicon carbide, etc. A particularly preferred and commercially available material of construction for operations below about 2,000°F. (1,093°C.) is cordierite, which is an alumina-magnesia-silica material. For applications involving operations above 2,000°F. (1,093°C) an alumina-silica-titania material is preferred. Honeycomb monoTithic supports are commercially available in various sizes and configurations. Typically, the monolithic carrier would comprise, e.g., a cordierite member of generally cylindrical configuration (either round or oval in cross section) and having a plurality of parallel gas flow passages or regular polygonal cross section extending therethrough. The gas flow passages are typically sized to provide from about 50 to 1,200, preferably 200 to 600, gas flow channels per square inch of face area.

Various honeycombed (reticulated) ceramic structures are described in the art: U.S. Pat. No. 4,251,239 discloses fluted filter of porous ceramic having increased surface area; U.S. Pat. No. 4,568,595 discloses honeycombed ceramic 20 foams with a surface having a ceramic sintered coating closing off the cells; U.S. Pat. No. 3,900,646 discloses ceramic foam with a nickel coating followed by platinum deposited in a syapor process; U.S. Pat. No. 3,957,685 bds still discloses nickel or spalladium coated on a negative image 25 ceramical metal/ceramicaloro metal foam: U.S. copper deposited in two layers with the second layer ssem or reinforced with aluminum, magnesium or zinc; U.S. Pat. No. bobulous 24,863,7120 discloses an negative image honeycombed 30 (reticulated) foam coated with cobalt, nickel or molybdenum coating; U.S. Pat. No. 4,308,233 discloses a reticulated to most sceramic foam having an mactivated alumina coating and a . Brown to go noble metal coating queeful as an exhaust gas catalyst; U.S. dove plainePat. Not 4,253,302 discloses, a foamed ceramic containing 35 35 platinum/rhodium catalyst for exhaust gas catalyst; and nul w in u.s. Pat. Non 4,088,607 discloses a ceramic foam having an was no coated by a noble metal containing composition such as zinc oxide, platinum and

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palladium.

The foam structure is characterized by the number of pores per linear inch and typical foams are produced with 10 to 100 pores per linear inch. The ceramic supports employed in the present invention are generally of the type disclosed in U.S. Pat. No. 4,810,685 using the appropriate material for the matrix and are generally referred to in the art and herein as "monoliths".

Generally any organic liquid in which the metal salt is soluble may be used to deposit metals on to monolith supports. The metals may also be deposited from aqueous solutions using the water soluble salts.

Generally from 0.5% to 20 wt % of the metal will be deposited on the monolith (based on the weight of monolith):

A suitable high surface area refractory metal oxide support layer may be deposited on the carrier to serve as a support upon which finely dispersed catalytic metal may be distended. As is known in the art, generally, oxides of one or more of the metals of Groups II, III, and IV of the Periodic Table of Elements having atomic numbers not greater than 40 are satisfactory as the support layer. Preferred high surface area support coatings are alumina, beryllia, zirconia, baria-alumina, magnesia, silica, and combinations of two or more of the foregoing.

The most preferred support coating is alumina, most preferably a stabilized, high-surface area transition alumina. One or more stabilizers such as rare earth metal oxides and/or alkaline earth metal oxides may be included in the transition alumina (usually in amounts comprising from 20 to 10 weight percent.

The metal monolith may be prepared as metal foam or sintered particles of metal at high temperature. Monolithic supports may also be made from materials such as nickel or stainless steel by placing a flat and a corrugated metal sheet, one over the other, and rolling the stacked sheets into a tubukar configuration about an axis parallel to the corrugations, to provide a cylindrical-

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shaped body having a plurality of fine, parallel gas flow passages extending therethrough.

The transition metals useful in the present invention are selected from the groups consisting of Fe, Os, Co, Rh, Ir, Ni Cu, Pd, Pt and mixtures thereof, with Fe, Co, Ni or Cu forming a preferred grouping, more preferably Ni and more preferably as a solid nickel monolith. Nickel supported on alumina monoliths was not found to be useful in the present process.

The noble metals useful in the present invention are selected from the group consisting of Ru, Rh, Pd, Pt and mixtures thereof preferably Pt or Pd.

For Figs. 1-10 runs were carried out in a quartz tube continuous reactor with 18 mm diameter. Identical α -Al₂O₃ monoliths without metal were positioned before and after the catalyst to reduce radiation losses. The catalyst and radiation shields were sealed in the quartz reactor by silica-alumina cloth. The temperatures of the front and back sides of the catalyst were measured with Pt-Pt/Rh thermocouples placed between the catalyst and the radiation shields. Rh and Pt catalysts were ignited with a Bunsen burner at a CH_4/O_2 ratio of 1.8. The fresh Ni spheres were more difficult to ignite, and therefore NH₃ was added to the feed to lower ignition temperature. After ignition, the reactor was insulated by wrapping it in high temperature insulation.

Rh and Pt catalysts were prepared by impregnating α -Al₂O₃ foam monoliths (18 mm diameter by 10 mm long) with concentrated metal salt solutions (rhodium chloride in acetone or chloroplatinic acid in H₂O) which were added dropwise to the monolith and allowed to dry overnight. The samples were further dried at 390 K for 2 hours and then calcined at 870° K for 2 hours in He and reduced at the same temperature in 10% H₂/Ar for 7 hrs. However, spheres of sintered Ni activated over a period of 6 hours (at normal reaction conditions) catalyzed an oxidation reaction in which the conversion and selectivities steadily increased. Therefore pure sintered Ni metal spheres (Alfa,

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-40 mesh, deposition grade, 99.9%) packed to 1 cm deep were used directly. Similar results were obtained with foamed nickel monolith.

Flow rates of high purity O21 N21 CO2 and CH4 were controlled by mass flow controllers, with an accuracy of 15 14 1/±0.01 standard liters per minute, (slpm)... experiments with no CO2 or H2O added to the feed were Low les or conducted at 4. slpm, CH₄/O₂= 1.8. and 35% dilution, which is optimal fuel/oxygen ratio for syngas production. of the ${\tt CO}_2$ results presented here, the ${\tt N}_2$ was replaced by CO2 to maintain a constant contact time within the The exit temperature of the catalyst was held catalyst. constant by preheating the feed gases. Runs were repeated where CO2 was added to a constant flow of CH4, O2 and N2. 150 G These results are nearly identical and show the same trends as the data presented here and are omitted. For runs in which steam was added to the feed, water was supplied by a syringe pump through a two stage vaporizer and a back de the pressure regulator to eliminate system pulsing. 102000 results reported, H20 was added to a constant flow of CH4, $(O_2, and)N_2$. Say they there if the (

chromatograph with a thermal conductivity detector and integrated by an on-line computer. The detected reaction products were H₂, CO, CO₂ and H₂O on all catalysts, and over Pt up to 1% C₂ products were observed. CH₄ conversion products selectivities were calculated as described previously. For CO₂ addition, the selectivities to CO and CO₂ are not reported (the only carbon containing product selectivities are not reported for H₂O addition. In all runs the carbon and hydrogen balances closed with an accuracy better than 97%.

Figs. 1-3 show the conversions of CH_4 and selectivities to H_2 with CO_2 addition on Ni, Rh and Pt catalysts. The reaction temperatures on Ni, Rh and Pt catalysts were held constant with CO_2 addition, 1240°, 1250° and 1500° K, respectively. The equilibrium predictions of CH_4

conversion and H_2 selectivities at these temperatures are shown by the dashed lines. Since Pt produces a lower H_2/H_2O ratio, its temperature is $\approx 250^{\circ}C$ hotter than Ni or Rh.

The CH₄ conversions were nearly constant up to 30% CO₂ addition on all three catalysts, but significantly lower than the equilibrium CH₄ conversions which is nearly 100%. These results imply that little CO₂ reforming of CH₄ occurs at contact times of ≈5 ms and ≈1300°K. Fig. 1 also shows 10 that the conversion of CH₄ is higher on Rh (80%) than on Ni (72%) with Pt (54%) being much less active.

The addition of CO₂ decreases the selectivity to H₂ over all three catalysts, but to very different amounts. With 24% CO₂ addition on Ni, the H₂ selectivity decreases from 15 80% to 55%, on Rh it decreases from 84% to 67%, and on Pt it decreases from 59% to 41%. The H₂ selectivities are roughly parallel to, but lower than, the calculated equilibrium H₂ selectivities. These results indicate that CO₂ reacts with H₂ in the reverse water-gas shift reaction to a greater extent on Nicand least on Pt.

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The results of H₂O addition with a constant flow of CH₄,
O₂, and N₂ over Ni, Rh oraPt are shown in Figs. 4-6. In
these runs, the temperature was kept constant at the
adiabatic temperature in the absence of H₂O by preheating
the gas stream. Because of preheat limitations,
maintaining a constant temperature on the Rt catalyst at
greater than 12 Vol% H₂O feed was difficult and therefore

Figs. 4-6 show that the CH47 conversions were nearly constant over the entire range of H20 addition on all three catalysts, indicating that negligible steam reforming of CH4 is occurring under these conditions.

As H₂O ds added to the feed the selectivity to CO decreases on Ni and Rh but remains essentially constant over Pt. On Ni, the selectivity to CO decreases from 88% to 50% While on Rh the selectivity only drops from 94% to 87%. The H₂O seems to be reacting with CO by the water-gas

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shift reaction over Ni and Rh but not over Pt.

The present results exhibited no catalyst deactivation on unsupported Ni and on supported Pt and Rh for over 100 hours of operation at atmospheric pressure. The present 5 data also show that as the concentration of CO2 in the feed increases, the conversion of CHA remains constant. Clearly shows that CO2 reforming is not occurring significantly at temperatures near 1300° K, at contact times near 1 ms and in the presence of oxygen.

Although these results may seem contradictory previous literature, three distinct factors differentiate this work from previous results of First, the present Freactor runs adiabatically and autothermally, therefore the reaction temperature is controlled by the catalyst selectivity and the temperatures typically run 250 to 500° hotter than typical CO2 reforming runs. Second, the contact time overwithe catalystem in our runs, is approximately 1 ms, corresponding to a gas hourly space velocity GHSV of 1x105h7, which is approximately a factor of 10 higher 2than most previous CO20 reforming work. Finally, the presence of O2 may inhibit CO2 reforming reactions. The CH4 dis probably preacting with 02 gvery rapidly, and the extremely short contact times prevent the remaining CH4 from reacting with CO2.

25 25 Choudhary et al, ... anolistic catal selecti. 32 (1995) = 391-396 sover NiQ-CaO catalyst at os space velocities of up to 5x105cm3 g-1 h-1 and temperatures enalested up to 1900 C. For CO2 reforming in the absence of O2, the NiO-CaO catalyst coked very rapidly but when 02 is added Talke30 "The feed, the catalyst showed no deactivation due to with the coke formation. by They sused their results to show that the 30 addition of 05 overcomes the endothermic limitations of CO2 reforming by initially combusting part of the CH4. energy released by combustion can then be immediately used to drive the reforming reactions: They confirmed this by 33 Jan Calculating other ratio cof CO2 reforming to exidative of the Conversion of CH4 based on the chemistry being controlled in wars, by three reactions of partial oxidation of CH4 with O2 to

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syngas, 2) complete combustion of CH4, 3) and CO2 reforming grand transportation and landing and the le of CHA.

Although, this may be a valid explanation of the product distribution in these experiments, it is believed that present results are better described by the direct oxidation of CHA to syngas followed by water-gas shift. Figs. 1-3 show that as CO2 is added to the feed, the conversion of CH4 does not change when the outlet temperature is held constant, for example by preheating the feed, indicating that co, reforming of CH4 is probably not active at these contact times and temperatures. However, as CO2 is added the selectivity to H2 decreases over all three metals. This implies that CO2 is reacting with H2 in the reverse water-gas shift.

The present results show that the water-gas shift reaction being affected by sthewaddition of H2O, particularly over the Nio monolith. The H2O seems to be reacting with co in the forward water-gas shift to form CO2 and H2. The combination of the CO2 and H2O addition results strongly confirm the activity of the water-gas shift reaction and its reverse at >1000° Kland 1 ms contact time.

As with CO2 reforming, steam reforming seems to be inactive over Ni, Rh or Pt at high temperatures and millisecond contact times. This is probably because the CH4 preferentially reacts with 02 first, and the remaining CH4 does not react with the H20 at these extremely short contact times. This is expected from the relatively low sticking coefficients of CO24 and H20 compared to O2.

The present results show that addition of CO_2 or H_2O to the feed stream affects the selectivity to H2; and CO but leaves the conversion of CH4 umaffected. The short contact times, 10 to 10 ms, do not allow significant CO2 or steam reforming on any of the three metals, counds the dominant reaction is the direct oxidation of CH4 to CO and H2 22

Since the H2 selectivity decreases with the addition of CO2 (or CO selectivity decreases with the addition of H2O), the water-gas shift reaction or its reverse must be active

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at present reaction temperatures and contact times.

Therefore, it is believed that the direct partial oxidation of CH₄ to syngas followed by the water-gas shift reaction are the dominant reactions in the present synvention.

Figs. 1-6 show that at contact times near 1 ms, the direct partial oxidation of methane to syngas appears to be followed by water-gas shift or its reverse. Because the water-gas shift reaction is active at 1300° K and 1 ms, the product selectivities can be tuned from high CO to high H₂ content by introducing CO₂ or H₂O respectively into the feed stream.

The negligible steam or CO₂ reforming occurs in the presence of O₂ over unsupported Ni and α-Al₂O₃ supported Rh and Pt catalysts at contact times of 1-10 ms and 1300 K. However, the water-gas shift reaction and the reverse shift reaction are active over Ni and Rh, with Pt being less active. Over Ni, the CO/CO₂ ratio decreases to 1 at 30% H₂O addition, which tranlates to a H₂/CO ratio of 5. This demonstrates the addition of H₂O or CO₂ can be used to manipulate the ratio of H₂/CO in the product stream.

In Fig. 11 an apparatus with a single transition metal acceptance of the arrow into tubular reactor 10. H₂O (as 25 25 25 team) for CO₂ enters via line 12 into a bead packing 22. The tubular reactor 10 is wrapped with heating tape and the drops catalysts 14 is sandwiched between two heat shields 16 and woll insulated by 18.

In Fig. 12 an apparatus with a multi-catalyst zone is shown in tubular reactor 100. Hydrocarbon/O2 feed enters 100 and 108 in the direction of the arrow to flow through the catalyst of and other mixture contacts noble metal catalyst 114 then transition metal (preferred group) 115. The two catalyst monoliths are associated with heat shields 116 and the insulated by 118. The front portion of the reactor

containing the beads is wrapped with heating tape 120.

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The hydrocarbon/O₂ feed 208 enters the tubular reactor 200 and passes through noble metal monolith catalyst 214 which is sandwiched between heat shields 216. That section is insulated by 218. The reaction has been initiated by igniting the catalyst 214. The product from this partial oxidation passes into the bead packing 222 where it mixes with H₂O (steam) or CO₂ entering via 212. The mixture proceeds through transition metal (preferred group) monolith catalyst 215. Heating tape is provided to maintain the temperature from the exothermic partial oxidation.

It has been found that unsupported metal monoliths such as nickel monolith is advantageous over ceramic supported metals. Because of the high thermal conductivity of Ni, the monolith is a better thermostat and higher flow rates can be achieved with similar selectivities and conversions. By using unsupported Ni, the possibility of nickelaluminate formation and therefore a deactivation route is eliminated. These two advantages combined allow for operation over wider range of inlet conditions.

The multiple catalyst bed design takes advantage of a more selective catalyst for synthesis gas formation. In this design the first bed is a supported noble metal catalyst, preferably Rh. This catalyst is supported by a reticulated or extruded ceramic structure and the weight loadings of catalyst can range from 0.1% to in excess of 10%. Steam or CO₂ is injected immediately following the first catalyst bed prior to entering the second bed. The second bed requires energy input to maintain reaction. This catalyst can be a noble metal, transition metal or metal oxide, like Ni, Fe, or Cu.

To initiate reaction, the feed gases are introduced to the reactor at the desired flow rate and composition. Energy is added to the catalyst either electrically or thermally until the catalyst ignites. After catalyst ignition the energy source is removed, and preheat is the only external energy source required to maintain reaction temperatures between 800° and 1000°C. For the multi-bed

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reactor, the second and beyond stages may need to be maintained at temperature either by external heating or from the energy released in the first stage.

Gas hourly space velocities -10^5h^{-1} have been used for the process. Inlet compositions of fuel and oxygen were taken from literature sources for the optimal production of synthesis gas. Natural gas was simulated by CH₄, the dominant component of natural gas. The inlet compositions ranged from 64% fuel, 36% oxidant to 35% fuel, 20% oxidant

The introduction of H₂O into the feed stream results in a decrease in the CO/CO₂ ratio. With no H₂O the CO:CO₂ ratio is 8:1, but with 40% H₂O feed the ratio is reduced to 1.3:1 when using an unsupported bed of nickel. Adding CO₂ instead of H₂O results in a decrease in the H₂:H₂O ratio from 9:1 to 1:1 again over a nickel catalyst. Similar results have been demonstrated over Rh and Pt catalysts.

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The invention claimed is:

- 1. A process for the partial oxidation of hydrocarbons by feeding a stream containing hydrocarbon, oxygen and H₂O or CO₂ through a catalyst zone containing a catalytically effective amount of at least one transition metal monolith catalyst under partial oxidation conditions wherein said transition metal is selected from the group consisting of Fe, Ru, Os, Co, Rh; cIr, Ni, Cu; Pd, and Pt.
- The process according to claim it wherein the transition metal is Ni or.Rh.
- 3. The process according to claim 1 wherein said catalyst comprises a nickel metal monolith.
 - 4. The process according to claim 1 wherein said catalyst zone contains one transition metal catalyst.
 - 5. The process according to claim 4 wherein the transition metal is Nicor Rhams 14
 - 6. The process according to claim 1 wherein H2O is fed with said hydrocarbon and soxygen and sort en
- 7. The process according to claim 1 wherein CO2 is fed with said hydrocarbon and oxygen and oxygen.
- 8. The process according to claim lawherein at least said hydrocarbon and said oxygen are contacted with a noble metal monolith catalyst under partial oxidation conditions prior to contacting said transition metal monolith catalyst wherein said mobble metal cisc selected from the group consisting of Ru, Rh, Pd, Ir, and Pt 13 10 2000
- of Caracata With said hydrocarbon sando exygen to contact said noble metal catalyst.
 - 10. The process according to claim 8 wherein ${\rm CO_2}$ is fed with said hydrocarbon and oxygen to contact said noble metal catalyst.
 - 11. The process according to claim 8 wherein said $\rm H_2O$ is fed to said stream after said noble metal monolith catalyst and prior to said transition metal monolith catalyst.
 - 12. The process according to claim 8 wherein said ${\rm CO}_2$ is fed to said stream after said noble metal monolith

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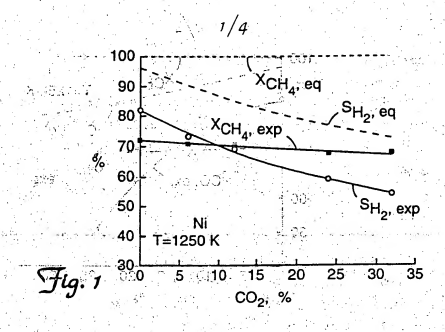
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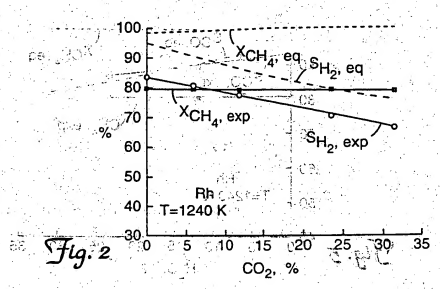
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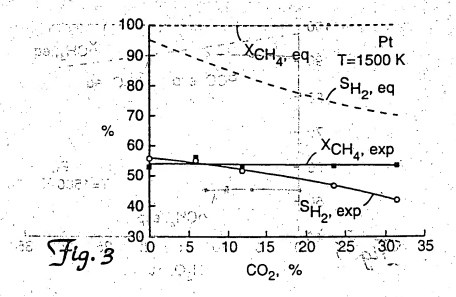
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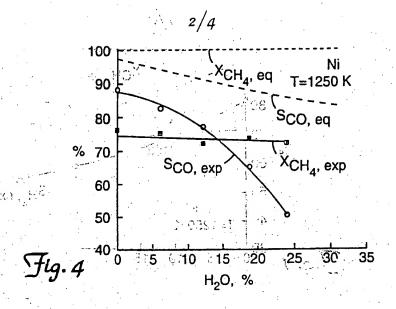
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catalyst and prior to assaids transition metal monolith
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13. The process according to claim 1, 4, 6, 7, 8, 9,
10, 11, or 12, wherein said transition metal is selected
from the group consisting of Fe, Ru, Os, Co, Rh, Ir, Ni,
                         nita in this and Cu.
10 gras tand 1440 The process saccording to claim 13 wherein said
       transition metal is Fe, Cu, for Ni.
and the 15. The process raccording to claim 14 wherein said
        transition metal is Cu. Lor of httlsdam noinely
     16. The process according to claim 14 wherein said
         transition ametal isoNi: on a combugate boy. . . .
         17. The process according to claim 8, 9, 10, 11, or 12,
      wherein said noble metalm is selected from the group
    tisted consisting of Pd and Ptune Responsibility
            18. The process according to claim 17 wherein said
      noble metal is Pt. paibarnes a enough and
            19. The process according to claim 17. wherein said
          noble metal is Pd. #15; nobbe waste ig add - 4.0
            20. The process according to claim 3 wherein said
          nickel monolith comprises a foamout at .3
said
nickel monolith comprises sintered nickel particles.
ray laura mosio 22. In the process according to claim 1 wherein the feed
guous gas to the reaction is preheated to a temperature in the
          range of 25-450 %C. (3) (44 ) 28 18 18 pails and
bet at 03H 40.2123: 3 The oprocessioaccording to claim 22 wherein the
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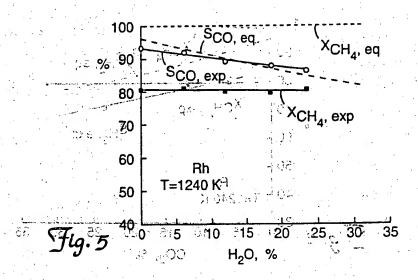
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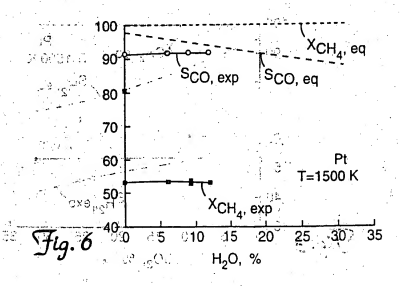


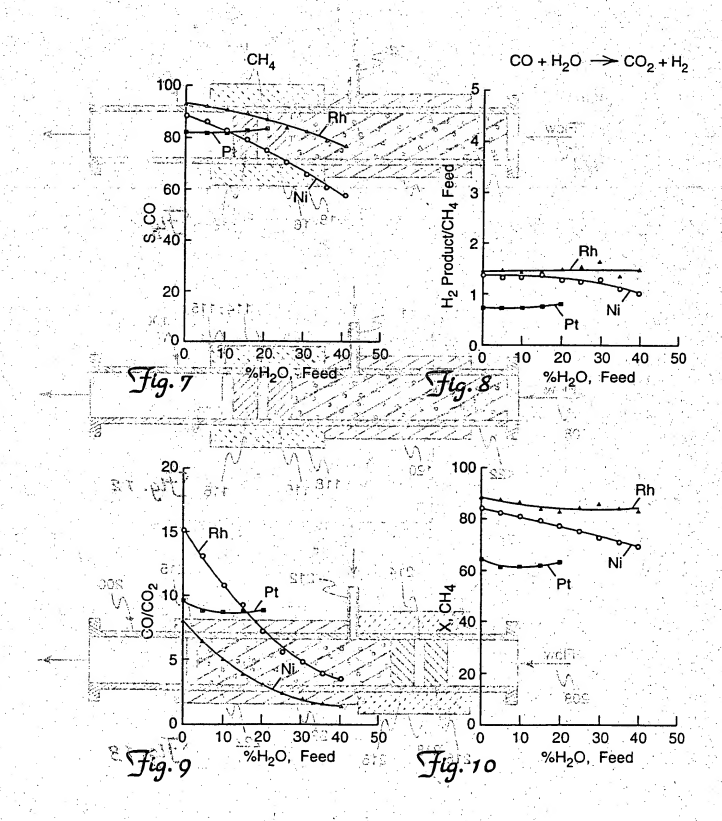


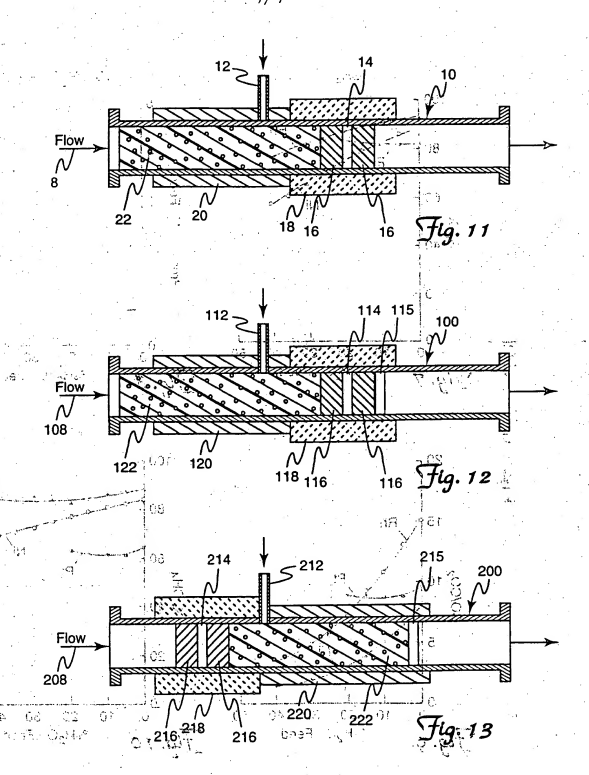












INTERNATIONAL SEARCH REPORT

ational Application No PCT/US 99/00629

A. CLASSIFICATION OF SUBJECT MATTER IPC. 6 ... C01B3/38

According to International Patent Classification (IPC) or to both national classification

Minimum documentation searched (classification system followed by classification symbols)

CO1B IPC 6

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical search terms used)

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Further documents are listed in the	continuation of box C.

Patent family members are listed in annex.

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- document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.

Date of mailing of the international search report

document member of the same patent family

Date of the actual completion of the international search

14 April 1999

Name and mailing address of the ISA

European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl. Fax: (+31-70) 340-3016

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	tinuation) DOCUMENTS CONSIDERED TO BE RELEVANT	The Control of the Co
Catego	ry ° Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	P. M. TORNIAINEN ET AL.: "Comparison of monolith-supported metals for the direct oxidation of methane to syngas"	1-6,13, 14,16,20
	JOURNAL OF CATALYSIS, vol. 146, 1994, pages 1-10, XP000770089 USA see page 6: effect of steam	
X	V. R. CHOUDHARY ET AL.: "Energy efficient methane-to-syngas conversion with low H2/CO ratio by simultaneous catalytic reactions of methane with carbon dioxide and oxygen" CATALYSIS LETTERS, vol- 32, 1995, pages 391-396, -XP002099617 BASEL; SWITZERLAND	1-5,7, 13,14,16
	cited in the application see the whole document	
X	see examples	1-5,7, 13,14,16
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INTERNATIONAL SEARCH REPORT

information on patent family members

Invational Application No PCT/US 99/00629

Carrier Contraction	Patent document cited in search report	Publication date	Patent family () member(s)	Publication date
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